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(54) Titre: METHODE ET MEDICAMENT DESTINES A INHIBER L'EXPRESSION D'UN GENE DONNE (54) Title: METHOD AND MEDICAMENT FOR INHIBITING THE EXPRESSION OF A GIVEN GENE

(57) Abrégé/Abstract:
The invention relates to a medicament containing at least one double-stranded oligoribonucleotide (dsRNA) designed to inhibit the expression of a target gene. According to the invention, one strand of the dsRNA is at least in part complementary to the target gene.





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- (57) Abstract

The invention relates to a medicament containing at least one double-stranded oligoribonucleotide (dsRNA) designed to inhibit the expression of a target gene. According to the invention, one strand of the dsRNA is at least in part complementary to the target gene.

(57) Zusammenfassung

Die Erfindung betrifft ein Medikament mit mindestens einem Oligoribonukleotid mit doppelsträniger Struktur (dsRNA) zur Hemmung der Expression eines Zielgens, wobei ein Strang der dsRNA zumindest abschnittsweise komplementär zum Zielgen ist.

(57) Abstract

The invention rolates to a medicament containing at least one double-stranded eligoribonucleotide (daRNA) designed to inhibit the expression of a target gene. According to the invention, one strand of the daRNA is at least in part complementary to the target gene.

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Method and medicament for inhibiting the expression of a given gene

The invention relates to methods in accordance with the 5 preambles of claims 1 and 2. It furthermore relates to double-stranded of medicament use and to oligoribonucleotides and to a vector encoding them.

Such a method is known from WO 99/32619, which was unpublished at the priority date of the present invention. The known process aims at inhibiting the expression of genes in cells of invertebrates. To this oligoribonucleotide double-stranded the end. exhibit a sequence which is identical with the target gene and which has a length of at least 50 bases. To 15 achieve efficient inhibition the lidentical sequence must be 300 to 12000 base pairs in length. Such an oligoribonucleotide is complicated to prepare.

DE 196 31 919 C2 describes with RNA an antisense 20 specific secondary structures, the antisense RNA being present in the form of a vector encoding it. The antisense RNA takes the form of an RNA molecule which is complementary to regions of the mRNA. Inhibition of the gene expression is caused by binding to these regions. This inhibition can be employed in particular for the diagnosis and/or therapy of diseases, for example tumor diseases or viral infections. - The disadvantage is that the antisense RNA must be introduced into the cell in an amount which is at least as high as the amount of the mRNA. The known antisense methods are not particularly effective.

medicament comprising US 5,712,257 discloses mismatched double-stranded RNA (dsRNA) and bioactive 35 mismatched fragments of dsRNA in the form of a ternary complex together with a surfactant. The dsRNA used for this purpose consists of synthetic nucleic acid single strands without defined base sequence. The single

"non-Watson-Crick" base pairing, giving rise to mismatched double strands. The known dsRNA is used to inhibit the amplification of retroviruses such as HIV. Amplification of the virus can be inhibited when non-sequence-specific dsRNA is introduced into the cells. This leads to the induction of interferon, which is intended to inhibit viral amplification. The inhibitory effect, or the activity, of this method is poor.

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It is known from Fire, A. et al., NATURE, Vol. 391, pp. 806 that dsRNA whose one strand is complementary in segments to a nematode gene to be inhibited inhibits the expression of this gene highly efficiently. It is believed that the particular activity of the dsRNA used in nematode cells is not due to the antisense principle but possibly on catalytic properties of the dsRNA, or enzymes induced by it. - Nothing is mentioned in this paper on the activity of specific dsRNA with regard to inhibiting the gene expression, in particular in mammalian and human cells.

The object of the present invention is to do away with the disadvantages of the prior art. In particular, it is intended to provide as effective as possible a method, medicament or use for the preparation of a medicament, which method, medicament or use is capable of causing particularly effective inhibition of the expression of a given target gene.

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This object is achieved by the features of claims 1, 2, 37, 38 and 74 and 75. Advantageous embodiments can be seen from claims 3 to 36, 39 to 73 and 76 to 112.

In accordance with the method-oriented inventions, it is provided in each case that the region I which is complementary to the target gene exhibits not more than 49 successive nucleotide pairs.

Provided in accordance with the invention are an oligoribonucleotide or a vector encoding therefor. At least segments of the oligoribonucleotide exhibit a defined nucleotide sequence. The defined segment may be limited to the complementary region I. However, it is also possible that all of the double-stranded oligoribonucleotide exhibits a defined nucleotide sequence.

it emerged that Surprisingly, has effective an 10 inhibition of the expression of the target gene can be achieved even when the complementary region I is not more than 49 base pairs in length. The procedure of providing such oligoribonucleotides is less 15 complicated.

particular, with In dsRNA a length over of pairs induces 50 nucleotide certain cellular mechanisms, for example the dsRNA-dependent protein kinase or the 2-5A system, in mammalian and human 20 leads to the disappearance of cells. This interference effect mediated by the dsRNA which exhibits a defined sequence. As a consequence, protein biosynthesis in the cell is blocked. The present 25 invention overcomes this disadvantage in particular.

Furthermore, the uptake of dsRNA with short chain lengths into the cell or into the nucleus is facilitated markedly over longer-chain dsRNAs.

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It has proved advantageous for the dsRNA or the vector to be present packaged into micellar structures, preferably in liposomes. The dsRNA or the vector can likewise be enclosed in viral natural capsids or in chemically or enzymatically produced artificial capsids or structures derived therefrom. - The abovementioned features make it possible to introduce the dsRNA or the vector into given target cells.

In a further aspect, the dsRNA has 10 to 1000, preferably 15 to 49, base pairs. Thus, the dsRNA can be longer than the region I, which is complementary to the target gene. The complementary region I can be located at the terminus or inserted into the dsRNA. Such dsRNA or a vector provided for coding the same can be produced synthetically or enzymatically by customary methods.

10 The gene to be inhibited is expediently expressed in eukaryotic cells. The target gene can be selected from the following group: oncogene, cytokin gene, Id protein gene, developmental gene, prion gene. It can also be expressed in pathogenic organisms, preferably in plasmodia. It can be part of a virus or viroid which is preferably pathogenic to humans. - The method proposed makes it possible to produce compositions for the therapy of genetically determined diseases, for example cancer, viral diseases or Alzheimer's disease.

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The virus or viroid can also be a virus or viroid which is pathogenic to animals or plant-pathogenic. In this case, the method according to the invention also permits the provision of compositions for treating animal or plant diseases.

In a further aspect, segments of the dsRNA are designed as double-stranded. A region II which is complementary within the double-stranded structure is formed by two separate RNA single strands or by autocomplementary regions of a topologically closed RNA single strand which is preferably in circular form.

The ends of the dsRNA can be modified to counteract degradation in the cell or dissociation into the single strands. Dissociation takes place in particular when low concentrations or short chain lengths are used. To inhibit dissociation in a particularly effective fashion, the cohesion of the complementary region II,

which is caused by the nucleotide pairs, can be increased by at least one, preferably two, further chemical linkage(s). - A dsRNA according to the invention whose dissociation is reduced exhibits greater stability to enzymatic and chemical degradation in the cell or in the organism.

The complementary region II can be formed by autocomplementary regions of an RNA hairpin loop, in particular when using a vector according to the invention. To afford protection from degradation, it is expedient for the nucleotides to be chemically modified in the loop region between the double-stranded structure.

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The chemical linkage is expediently formed by a covalent or ionic bond, a hydrogen bond, hydrophobic interactions, preferably van-der-Waals or stacking interactions, or by metal-ion coordination. In an especially advantageous aspect, it can be formed at at least one, preferably both, end(s) of the complementary region II.

It has furthermore proved to be advantageous for the chemical linkage to be formed by one or more linkage linkage the groups preferably poly(oxyphosphinicooxy-1,3-propanediol) and/or polyechylene glycol chains The chemical linkage can also be formed by purine analogs used in place of purines in the complementary regions II. It is also advantageous 30 for the chemical linkage to be formed by azabenzene units introduced into the complementary regions II. Moreover, it can be formed by branched nucleotide used in place of nucleotides analogs the complementary regions II. 35

It has proved expedient to use at least one of the following groups for generating the chemical linkage: methylene blue; bifunctional groups, preferably

bis(2-chloroethyl)amine; N-acetyl-N'-(p-glyoxylbenzoyl) cystamine; 4-thiouracil; psoralene. The linkage chemical furthermore can be formed thiophosphoryl groups provided at the ends of double-stranded region. The chemical linkage at ends of the double-stranded region is preferably formed by triple-helix bonds.

The chemical linkage can expediently be induced by ultraviolet light.

The micleotides of the dsRNA can be modified This counteracts the activation, in the cell, of a doublestranded-RNA-dependent protein kinase, And the second of the second o Advantageously, at least one 2'-hydroxyl group of the 15 nucleotides of the dsRNA in the complementary region II is replaced by a chemical group, preferably a 2'-amino. or a 2'-methyl group. At least one nucleotide in at least one strand of the complementary region II can also be a locked nucleotide with a sugar ring which is chemically modified, preferably by methylene bridge. Advantageously, several nucleotides are locked nucleotides.

25 A further especially advantageous embodiment provides that the dsRNA or the vector is bound to, associated with or surrounded by, at least one viral coat protein which originates from a virus, is derived therefrom or has been prepared synthetically. The coat protein can be derived from polyomavirus. The coat protein can 30 contain the polyomavirus virus protein 1 (VP1) and/or virus protein 2 (VP2). The use of such coat proteins is for example, known from, DE 196 18 797 A1, disclosure is herewith incorporated. The abovementioned features considerably facilitate the 35 introduction of the dsRNA or of the vector into the cell.

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When a capsid or capsid-type structure is formed from the coat protein, one side preferably faces the interior of the capsid or capsid-type structure. The construct formed is particularly stable.

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The dsRNA can be complementary to the primary or processed RNA transcript of the target gene. - The cell can be a vertebrate cell or a human cell.

10 At least two dsRNAs which differ from each other or at least one vector encoding them can be introduced into the cell, where at least segments of one strand of each dsRNA are complementary to in each case one of at least two different target genes. This makes it possible simultaneously to inhibit the expression of at least two different target genes. In order to suppress, in the cell, the expression of a double-stranded-RNA-dependent protein kinase, PKR, one of the target genes is advantageously the PKR gene. This allows effective suppression of the PKR activity in the cell.

The invention furthermore provides a medicament with at one oligoribonucleotide with double-stranded least structure (dsRNA) for inhibiting the expression of a given target gene, where one strand of the dsRNA has a region I where at least segments are complementary to the target gene. - Surprisingly, it has emerged that such a dsRNA is suitable as medicament for inhibiting the expression of a given gene in mammalian cells. In comparison with single-stranded the use of oligoribonucleotides, the inhibition is already caused at concentrations which are lower by at least one order of magnitude. The medicament according to the invention is highly effective. Lesser side effects can be expected.

The invention furthermore provides a medicament with at least one vector for coding at least one oligoribonucleotide with double-stranded structure

(dsRNA) for inhibiting the expression of a given target gene, where one strand of the dsRNA has a region I where at least segments are complementary to the target gene. - The medicament proposed exhibits the abovementioned advantages. By using a vector, in particular production costs can be reduced.

In a particularly advantageous embodiment, the complementary region I has not more than 49 successive nucleotide pairs. - Surprisingly, it has emerged that an effective inhibition of the expression of the target gene can be achieved even when the complementary region I is not more than 49 base pairs in length. The procedure of providing such oligoribonucleotides is less complicated.

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The invention furthermore provides a use of an oligoribonucleotide with double-stranded structure (dsRNA) for preparing a medicament for inhibiting the expression of a given target gene, where one strand of the dsRNA has a region I where at least segments are complementary to the target gene. - Surprisingly, such a dsRNA is suitable for preparing a medicament for inhibiting the expression of a given gene. Compared with the use of single-stranded oligoribonucleotides, the inhibition is already caused at concentrations which are lower by one order of magnitude when using dsRNA. The use according to the invention thus makes possible the preparation of particularly effective medicaments.

The invention furthermore provides the use of a vector for coding at least one oligoribonucleotide with double-stranded structure (dsRNA) for preparing a medicament for inhibiting the expression of a given target gene, where one strand of the dsRNA has a region I where at least segments are complementary to this target gene. - The use of a vector makes possible a particularly effective gene therapy.

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With regard to advantageous embodiments of the medicament and of the use, reference is made to the description of the above features.

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Use examples of the invention are illustrated in greater detail hereinbelow with reference to the figures, in which:

- 10 Fig. 1 shows the schematic representation of a plasmid for the in vitro transcription with T7- and SP6-polymerase,
- Fig. 2 shows RNA following electrophoresis on an 8% polyacrylamide gel and staining with ethidium bromide,
- Fig. 3 shows a representation of radioactive RNA transcripts following electrophoresis on an 8% polyacrylamide gel with 7 M urea by means of an instant imager, and
 - Figs. 4a e show Texas Red and YFP fluorescence in murine fibroblasts.

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Use example 1:

The inhibition of transcription was detected by means of sequence homologous dsRNA in an in vitro transcription system with a nuclear extract from human HeLa cells. The DNA template for this experiment was plasmid pCMV1200 which had been linearized by means of BamHI.

Generation of the template plasmids:

The plasmid shown in fig. 1 was constructed for use in the enzymatic synthesis of the dsRNA. To this end, a polymerase chain reaction (PCR) with the "positive control DNA" of the HelaScribe" Nuclear Extract in vitro transcription kit by Promega, Madison, USA, as

DNA template was first carried out. One of the primers used contained the sequence of an EcoRI cleavage site and of the T7 RNA polymerase promoter as shown in sequence listing No. 1. The other primer contained the sequence of a BamHI cleavage site and of the SP6 RNA polymerase promoter as shown in sequence listing No. 2. In addition, the two primers had, at the 3' ends, regions which were identical with or complementary to the DNA template. The PCR was carried out by means of the "Tag PCR Core Kits" by Qiagen, Hilden, Germany, 10 following the manufacturer's instructions. $MgCl_2$, in each case 200 μM dNTP, in each case 0.5 μM primer, 2.5 U Taq DNA polymerase and approximately 100 ng of "positive control DNA" were employed as template in PCR buffer in a volume of 100 μ l. After 15 initial denaturation of the template DNA by heating for 5 minutes at 94°C, amplification was carried out in 30 cycles of denaturation for in each case 60 seconds at 94°C, annealing for 60 seconds at 5°C below the melting point primers and calculated οf the 20 polymerization for 1.5-2 minutes at 72°C. After a final polymerization of 5 minutes at 72°C, 5 µl of the reaction were analyzed by agarose-gel electrophoresis. The length of the DNA fragment amplified thus was 400 base pairs, 340 base pairs corresponding to the 25 "positive control DNA". The PCR product was purified, after hydrolyzed with **ECORI** BamHI and, and repurification, employed in the ligation together with a pUC18 vector which had also been hydrolyzed by EcoRI and BamHI. E. coli XL1-blue was then transformed. The 30 plasmid obtained (pCMV5) carries a DNA fragment whose 5' end is flanked by the T7 promoter and whose 3' end is flanked by the SP6 promoter. By linearizing the plasmid with BamHI, it can be employed in vitro with the T7-RNA polymerase for the run-off transcription of a single-stranded RNA which is 340 nucleotides in length and shown in sequence listing No. 3. If the plasmid is linearized with EcoRI, it can be employed for the run-off transcription with SP6 RNA polymerase,

giving rise to the complementary strand. In accordance with the method outlined hereinabove, an RNA 23 nucleotides in length was also synthesized. To this end, a DNA shown in sequence listing No. 4 was ligated with the pUC18 vector via the *EcoRI* and *BamHI* cleavage sites.

Plasmid pCMV1200 was constructed as DNA template for the in-vitro transcription with HeLa nuclear extract. To this end, a 1 191 bp EcoRI/BamHI fragment of the 10 positive control DNA contained in the HeLaScribe® Nuclear Extract in vitro transcription kit was amplified by means of PCR. The amplified fragment encompasses the 828 bp "immediate early" CMV promoter and a 363 bp transcribable DNA fragment. 15 ligated to the vector pGEM-T via product was "T-overhang" ligation. A BamHI cleavage site is located at the 5' end of the fragment. The plasmid was linearized by hydrolysis with BamHI and used as template in the run-off transcription. 20

In-vitro transcription of the complementary single strands:

pCMV5 plasmid DNA was linearized with EcoRI or BamHI. It was used as DNA template for an in-vitro transcription of the complementary RNA single strands with SP6 and T7 RNA polymerase, respectively. The "Riboprobe in vitro Transcription" system by Promega, Madison, USA, was employed for this purpose. Following the manufacturer's instructions, 2 μ g of linearized plasmid DNA were incubated in 100 μ l of transcription buffer and 40 U T7 or SP6 RNA polymerase for 5-6 hours at 37°C. The DNA template was subsequently degraded by addition of 2.5 μ l of RNase-free DNase RQ1 and incubation for 30 minutes at 37°C. The transcription reaction was made up to 300 μ l with H₂O and purified by phenol extraction. The RNA was precipitated by addition of 150 μ l of 7 M ammonium acatate (sic) and 1 125 μ l of

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ethanol and stored at -65°C until used for the hybridization.

Generation of the RNA double strands:

- For the hybridization, 500 μ l of the single-stranded RNA which had been stored in ethanol and precipitated were spun down. The resulting pellet was dried and taken up in 30 μ l of PIPES buffer, pH 6.4 in the presence of 80% formamide, 400 mM NaCl and 1 mM EDTA.
- In each case 15 μ l of the complementary single strands were combined and heated for 10 minutes at 85°C. The reactions were subsequently incubated overnight at 50°C and cooled to room temperature.
- Only approximately equimolar amounts of the two single strands were employed in the hybridization. This is why the dsRNA preparations contained single-stranded RNA (ssRNA) as contaminant. In order to remove these ssRNA contaminants, the reactions were treated, after hybridization, with the single-strand-specific
 - ribonucleases bovine pancreatic RNase A and Aspergillus oryzae RNase Tl. RNase A is an endoribonuclease which is specific for pyrimidines. RNase Tl is an endoribonuclease which preferentially cleaves at the 3'

side of quanosines, dsRNA is no substrate for these

- ribonucleases. For the RNase treatment, the reactions in 300 μ l of Tris, pH 7.4, 300 mM NaCl and 5 mM EDTA were treated with 1.2 μ l of RNaseA at a concentration of 10 mg/ml and 2 μ l of RNaseT1 at a concentration of
- 30 290 μ g/ml. The reactions were incubated for 1.5 hours at 30°C. Thereupon, the RNases were denatured by addition of 5 μ l of proteinase K at a concentration of 20 mg/ml and 10 μ l of 20% SDS and incubation for 30 minutes at 37°C. The dsRNA was purified by phenol
- extraction and precipitated with ethanol. To verify the completeness of the RNase digestion, two control reactions were treated with ssRNA analogously to the hybridization reactions.

The dried pellet was taken up in 15 μ l of TE buffer, pH 6.5, and subjected to native polyacrylamide gel electrophoresis on an 8% gel. The acrylamide gel was subsequently stained in an ethidium bromide solution and washed in a water bath. Fig. 2 shows the RNA which had been visualized in a UV transilluminator. The sense RNA which had been applied to lane 1 and the antisense RNA which had been applied to lane 2 showed a different migration behavior under the chosen conditions than the dsRNA of the hybridization reaction which had been applied to lane 3. The RNase-treated sense RNA and antisense RNA which had been applied to lanes 4 and 5, respectively, produced no visible band. This shows that the single-stranded RNAs had been degraded completely. The RNase-treated dsRNA of the hybridization reaction which had been applied to lane 6 is resistant to RNase treatment. The band which migrates faster in the native gel in comparison with the dsRNA applied to lane 3 results from dsRNA which is free from ssRNA. addition to the dominant main band, weaker bands which migrate faster are observed after the RNase treatment.

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In-vitro transcription test with human nuclear extract: the HeLaScribe Nuclear Extract Using in vitro kit by Promega, transcription 25 Madison, USA, the transcription efficiency of the abovementioned DNA fragment which is present in plasmid pCMV1200 and homologous to the "positive control DNA" was determined in the presence of the dsRNA (dsRNA-CMV5) with sequence homology. Also, the effect of the dsRNA without 30 sequence homology, which corresponds to the yellow fluorescent protein (YFP) (dsRNA-YRP), gene studied. This dsRNA had been generated analogously to the dsRNA with sequence homology. The sequence of a strand of this dsRNA can be found in sequence listing 35 No. 5. Plasmid pCMV1200 was used as template for the run-off transcription. It carries the "immediate early" cytomegalovirus promoter which is recognized by the eukaryotic RNA polymerase II, and a transcribable DNA

fragment. Transcription was carried out by means of the HeLa nuclear extract, which contains all the proteins which are necessary for transcription. By addition of $[\cdot - ^{32}P]$ rGTP to the transcription reaction, radiolabeled transcript was obtained. The $[\cdot -3^{2}P]$ rGTP used had a specific activity of 400 Ci/mmol, 10 mCi/ml. $MgCl_2$, in each case 400 μM rATP, rCTP, rUTP, 16 μM rGTP, 0.4 μ M [\cdot -³²P]rGTP and depending on the experiment 1 fmol of linearized plasmid DNA and various amounts of in transcription buffer were employed per dsRNA 10 reaction. Each batch was made up to a volume of $8.5 \mu l$ with H2O. The reactions were mixed carefully. To start the transcription, 4 U HeLa nuclear extract in a volume of $4 \mu l$ were added and incubated for 60 minutes at 30°C. The reaction was stopped by addition of 87.5 μ l of quench mix which had been warmed to 30°C. To remove the proteins, the reactions were treated with 100 μ l of phenol/chloroform/isoamyl alcohol (25:24:1 saturated with TE buffer, pH 5.0, and the reactions mixed vigorously for 1 minute. For 20 were separation, the reactions were spun for approximately 1 minute at 12 000 rpm the and top phase was transferred into a fresh reaction vessel. Each reaction was treated with 250 μ l of ethanol. The reactions were mixed thoroughly and incubated for at least 15 minutes dry ice/methanol. To precipitate the RNA, reactions were spun for 20 minutes at 12 000 rpm and 40°C. The supernatant was discarded. The pellet was dried in vacuo for 15 minutes and resuspended in 10 μ l of H_2O . Each reaction was treated with 10 μ l of denaturing loading buffer. The free GTP was separated from the transcript formed by means of denaturing polyacrylamide gel electrophoresis on an 8% gel with 7 M urea. The RNA transcripts formed upon transcription with HeLa nuclear extract, in denaturing loading buffer, were heated for 10 minutes at 90°C and 10 μ l aliquots were applied immediately to the freshly washed pockets. The electrophoresis was run at 40 mA. The of the radioactive SSRNA formed upon amount

transcription was analyzed after electrophoresis with the aid of an Instant Imager.

Fig. 3 shows the radioactive RNA from a representative test, shown by means of the *Instant Imager*. Samples obtained from the following transcription reactions were applied:

Lane 1: without template DNA, without dsRNA;

10 Lane 1: 50 ng of template DNA, without dsRNA;

Lane 3: 50 ng of template DNA, 0.5 μ g of dsRNA YFP;

Lane 4: 50 ng of template DNA, 1.5 µg of dsRNA YFP;

Lane 5: 50 ng of template DNA, 3 μ g of dsRNA YFP;

Lane 6: 50 ng of template DNA, 5 μ g of dsRNA YFP;

15 Lane 7: without template DNA, 1.5 dsRNA YFP;

Lane 8: 50 ng of template DNA, without dsRNA;

Lane 9: 50 ng of template DNA, 0.5 μ g of dsRNA CMV5;

Lane 10: 50 ng of template DNA, 1.5 μ g of dsRNA CMV5;

Lane 11: 50 ng of template DNA, 3 µg of dsRNA CMV5;

20 Lane 12: 50 ng of template DNA, 5 μg of dsRNA CMV5;

It emerged that the amount of transcript was reduced markedly in the presence of dsRNA with sequence in comparison with the control reaction homology without dsRNA and with the reactions with dsRNA YFP without sequence homology. The positive control in lane 2 shows that radioactive transcript was formed upon the in-vitro transcription with HeLa nuclear extract. The reaction is used for comparison with the transcription reactions which had been incubated in the presence of 30 dsRNA. Lanes 3 to 6 show that the addition of nonsequentially-specific dsRNA YFP had no effect on the amount of transcript formed. Lanes 9 to 12 show that the addition of an amount of between 1.5 and 3 μg of sequentially-specific dsRNA CMV5 leads to a reduction 35 in the amount of transcript formed. In order to exclude that the effects observed are based not on the dsRNA but on any contamination which might have been carried along accidentally during the preparation of the dsRNA,

a further control was carried out. Single-stranded RNA was transcribed as described above and subsequently subjected to the RNase treatment. It was demonstrated by means of native polyacrylamide gel electrophoresis that the ssRNA had been degraded completely. This reaction was subjected to phenol extraction and ethanol precipitation and subsequently taken up in PE buffer, as were the hybridization reactions. This gave a sample which contained no RNA but had been treated with the same enzymes and buffers as the dsRNA. Lane 8 shows that the addition of this sample had no effect on transcription. The reduction of the transcript upon addition of sequence-specific dsRNA can therefore be ascribed unequivocally to the dsRNA itself. reduction of the amount of transcript of a gene in the presence of dsRNA in a human transcription system indicates an inhibition of the expression of the gene in question. This effect can be attributed to a novel mechanism caused by the dsRNA.

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Use example 2:

The test system used for these in-vivo experiments was the murine fibroblast cell line NIH3T3, ATCC CRL-1658. The YFP gene was introduced into the nuclei with the aid of microinjection. Expression of YFP was studied under the effect of simultaneously cotransfected dsRNA with sequence homology. This dsRNA YFP shows homology with the 5'-region of the YFP gene over a length of 315 bp. The nucleotide sequence of a strand of the sequence listing shown dsrna yrp is in fluorescence under the microscope Evaluation carried out 3 hours after injection with reference to the greenish-yellow fluorescence of the YFP formed.

35 Construction of the template plasmid, and preparation of the dsRNA:

A plasmid was constructed following the same principle as described in use example 1 to act as template for the production of the YFP dsRNA by means of T7 and SP6

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in-vitro transcription. Using the primer Eco_T7_YFP as shown in sequence listing No. 6 and Bam_SP6_YFP as shown in sequence listing No. 7, the desired gene fragment was amplified by PCR and used analogously to the above description for preparing the dsRNA. The dsRNA YFP obtained is identical to the dsRNA used in use example 1 as non-sequence-specific control.

A dsRNA linked chemically at the 3' end of the RNA as shown in sequence listing No. 8 to the 5' end of the complementary RNA via a C18 linker group was prepared (L-dsRNA). To this end, synthons modified by disulfide bridges were used. The 3'-terminal synthon is bound to the solid support via the 3' carbon with an aliphatic linker group via a disulfide bridge. In the 5'-terminal synthon of the complementary oligoribonucleotide which is complementary to the 3'-terminal synthon of the one oligoribonucleotide, the 5'-trityl protecting group is bound via a further aliphatic linker and a disulfide bridge. Following synthesis of the two single strands, removal of the protecting groups and hybridization of complementary oligoribonucleotides, the the thiol groups which form are brought into spatial vicinity. The single strands are linked to each other by oxidation via their aliphatic linkers and a disulfide bridge. This is followed by purification with the aid of HPLC.

Preparation of the cell cultures:

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The cells were incubated in DMEM supplemented with 4.5 g/l glucose, 10% fetal bovine serum in culture dishes at 37°C under a 7.5% CO₂ atmosphere and passaged before reaching confluence. The cells were detached with trypsin/EDTA. To prepare for microinjection, the cells were transferred into Petri dishes and incubated further until microcolonies formed.

Microinjection:

For the microinjection, the culture dishes were removed incubator for approximately 10 minutes. from the Approximately 50 nuclei were injected singly per reaction within a marked area using the AIS microinjection system from Carl Zeiss, Göttingen, Germany. The cells were subsequently incubated for three more hours. For the microinjection, borosilicate glass capillaries from Hilgenberg GmbH, Malsfeld, Germany, with a diameter of less than 0.5 μ m at the tip were prepared. The microinjection was carried out using a micromanipulator from Narishige Scientific Instrument Lab., Tokyo, Japan. The injection time was 0.8 seconds the pressure was approximately 100 hPa. transfection was carried out using the plasmid pCDNA 15 YFP, which contains an approximately 800 bp BamHI/EcoRI fragment with the YFP gene in vector pcDNA3. The samples injected into the nuclei contained 0.01 μ g/ μ l of pCDNA-YFP and Texas Red coupled to dextran-70000 in 14 mM NaCl, 3 mM KCl, 10 mM KPO₄ [sic], ph 7.5. 20 Approximately 100 pl of RNA with a concentration of 1 μ M or, in the case of the L-dsRNA, 375 μ M were additionally added.

The cells were studied under a fluorescence microscope with excitation with the light of the excitation wavelength of Texas Red, 568 nm, or of YFP, 488 nm. Individual cells were documented by means of a digital camers. Figures 4a-e show the result for NIH3T3 cells.

In the cells shown in Fig. 4a, sense-YFP-ssRNA has been injected, in Fig. 4b antisense-YFP-ssRNA, in Fig. 4c dsRNA-YFP, in Fig. 4d no RNA and in Fig. 4e L-dsRNA.

The field on the left shows in each case the fluorescence of cells with excitation at 568 nm. The fluorescence of the same cells at an excitation of 488 nm is seen on the right. The Texas Red fluorescence of all the cells shown demonstrates that the injection solution had been applied successfully into the nuclei

and that cells with successful hits were still alive after three hours. Dead cells no longer showed Texas Red fluorescence.

The right fields of each of figures 4a and 4b show that YFP expression was not visibly inhibited when the single-stranded RNA was injected into the nuclei. The right field of Fig. 4c shows cells whose longer detectable after fluorescence was no injection of dsRNA-YFP. Fig. 4d shows cells into which 10 no RNA had been injected, as control. The cell shown in fig. 4e shows YFP fluorescence which can no longer be detected owing to the injection of the L-dsRNA which shows regions with sequence homology to the YFP gene. This result demonstrates that even shorter dsRNAs can be used for specifically inhibiting gene expression in mammals when the double strands are stabilized by chemically linking the single strands.

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Sequence Listing

- <110> Kreutzer Dr., Roland
 Limmer Dr., Stephan
- <120> Method and medicament for inhibiting the expression of a given gene
- <130> 400968
- <140>
- <141>
- <150> 199 03 713.2
- <151> 1999-01-30
- <150> 199 56 568.6
- <151> 1999-11-24
- <160> 8
- <170> PatentIn Ver. 2.1
- <210> 1
- <211> 45
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- <213> Artificial Sequence
- <220>
- coription of the artificial sequence:
 EcoRI cleavage site, T7 RNA Polymerase
 promoter

<400> 1
gganttetaa tacgacteae tatagggega teagatetet agaag

45

<210> 2

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2

50 <211>

DNA <212>

Artificial Sequence <213>

<220>

Description of the artificial sequence: <223> cleavage site, SP6 RNA Polymerase BamHI promoter

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340 <211>

RNA <212>

Artificial Sequence <213>

<220>

Description of the artificial sequence: <223> RNA which corresponds to a sequence from the positive control DNA of the HeLa Nuclear vitro transcription kit Extract in Promega

<400> 3

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<210> 4

<211> 363

<212> DNA

Artificial Sequence <213>

<220>

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3

DNA which corresponds to a sequence from the positive control DNA of the HeLa Nuclear Extract in vitro transcription kit from Promega

<400> 4

geacegigta igaaatetaa caaigegite alegicatee teggeacegi caecetggal 120 geigtaggea taggetiggi taigeeggia eigeeggigee tetigeggia talegiceat 180 teegacagea tegecagica eiaigegig eigetagege taiaigegii gaigeaalii 240 eiaigegeae eegitetegg ageaetgice gaeegetiig geegeegeee agicetgeic 300 geitegetae tiggageeae taieggiae gegaleaigg eigaeeacacaee egicetgiig 360 ate

<210> 5

<211> 315

<212> RNA

<213> Artificial Sequence

<220>

<223> Description of the artificial sequence:
 Sequence from the YFP gene

<400> 5

auggugagea agggegagga geuguucace ggggugguge ecauceuggu egageuggae 60 ggegaeguaa aeggeeacaa guucagegug uceggegagg gegagggega ugecaceuae 120 ggeaageuga eecugaaguu eaueugeace aeeggeaage ugeceggugee euggeeeace 180 euegugaeea eccugaecua eggegugeag ugeuucagee geuaeeeega eeaeaugaag 240 eageaegaeu ucuucaague egeeaugeee gaaggeuaeg uceaggageg eaeeaucuuc 300 uucaaggaeg aegge

<210> 6

<211> 52

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of the artificial sequence:

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EcoRI cleavage site, T7 RNA Polymerase promoter, complementary region to the YFP gene

<400> 6
ggaattetan tacgseteae tatagggega atggtgagea agggegagga ge 52

<210> 7

<211> 53

<212> DNA

<213> Artificial Sequence

<220>

223> Description of the artificial sequence:
 BamHI cleavage site, SP6 RNA Polymerase
 promoter, complementary region to the YFP gene

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gggatecatt taggtgacac tatagantac geogtegtee ttganganga tgg 53

<210> 8

<211> 21

<212> RNA

<213> Artificial Sequence

<220>

Color of the artificial sequence:

RNA which corresponds to a sequence from the Strategy of the Strategy

<400> \$

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International Patent Application No. PCT/DE00/00244 of Dr Roland Kreutzer and Dr Stefan Limmer

New Patent Claims

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- 1. Method for inhibiting the expression of a given target gene in a cell in vitro, where an oligoribonucleotide with double-stranded structure (dsRNA) formed by two separate RNA single strands is introduced into the cell, where one strand of the dsRNA has a region which is complementary to the target gene,
- characterized in that
 the complementary region has less than 25
 successive nucleotide pairs.
 - Method according to claim 1, where the dsRNA is enclosed by micellar structures, preferably by liposomes.

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- 3. Method according to either of the preceding claims, where the dsRNA is enclosed by natural viral capsids or by chemically or enzymatically produced artificial capsids or structures derived therefrom.
- 4. Method according to one of the preceding claims, where the target gene is expressed in eukaryotic cells.

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5. Method according to one of the preceding claims, where the target gene is selected from the following group: oncogene, cytokin gene, Idprotein gene, development gene, prion gene.

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6. Method according to one of the preceding claims, where the target gene is expressed in pathogenic organisms, preferably in plasmodia.

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- 7. Method according to one of the preceding claims, where the target gene is part of a virus or viroid.
- 8. Method according to claim 7, where the virus is a virus or viroid which is pathogenic for humans.
- 9. Method according to claim 7, where the virus or viroid is a virus or viroid which is pathogenic for animals or phytopathogenic.
- 10. Method according to one of the preceding claims, where segments of the dsRNA are in double-stranded form.
- 11. Method according to one of the preceding claims, where the ends of the dsRNA are modified in order to counteract degradation in the cell or dissociation into the single strands.
- 12. Method according to one of the preceding claims, where the cohesion of the double-stranded structure, which is caused by the complementary nucleotide pairs, is increased by at least one, preferably two, further chemical linkage(s).
- 13. Method according to one of the preceding claims, where the chemical linkage is formed by a covalent or ionic bond, a hydrogen bond, hydrophobic interactions, preferably van-der-Waals or stacking interactions, or by metal-ion coordination.
- 14. Method according to one of the preceding claims,
 where the chemical linkage is generated at at
 least one, preferably both, ends of the doublestranded structure.

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- 15. Method according to one of the preceding claims, where the chemical linkage is formed by means of one or more compound groups, the compound groups preferably being poly(oxyphosphinicooxy-1,3-propanediol) and/or polyethylene glycol chains.
- 16. Method according to one of the preceding claims, where the chemical linkage is formed by purine analogs used in the double-stranded structure in place of purines.

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- 17. Method according to one of the preceding claims, where the chemical linkage is formed by azabenzene units introduced into the double-stranded structure.
- 18. Method according to one of the preceding claims, where the chemical linkage is formed by branched nucleotide analogs used in the double-stranded structure in place of nucleotides.
- 19. Method according to one of the preceding claims, where at least one of the following groups is used for generating the chemical linkage: methylene blue; bifunctional groups, preferably bis(2-chloroethyl)amine; N-acetyl-N'-(p-glyoxyl-benzoyl)cystamine; 4-thiouracil; psoralene.
- 30 20. Method according to one of the preceding claims, where the chemical linkage is formed by thiophosphoryl groups provided at the ends of the double-stranded structure.
- 35 21. Method according to one of the preceding claims, where the chemical linkage at the ends of the double-stranded structure is formed by triple-helix bonds.

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- 22. Method according to one of the preceding claims, where at least one 2'-hydroxyl group of the nucleotides of the dsRNA in the double-stranded structure is replaced by a chemical group, preferably a 2'-amino or a 2'-methyl group.
- 23. Method according to one of the preceding claims, where at least one nucleotide in at least one strand of the double-stranded structure is a locked nucleotide with a sugar ring which is chemically modified, preferably by a 2'-0, 4'-C-methylene bridge.
- 15 24. Method according to one of the preceding claims, where the dsRNA is bound to, associated with or surrounded by, at least one viral coat protein which originates from a virus, is derived therefrom or has been prepared synthetically.

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- 25. Method according to one of the preceding claims, where the coat protein is derived from polyomavirus.
- 25 26. Method according to one of the preceding claims, where the coat protein contains the polyomavirus virus protein 1 (VP1) and/or virus protein 2 (VP2).
- 30 27. Method according to one of the preceding claims, where, when a capsid or capsid-type structure is formed from the coat protein, one side faces the interior of the capsid or capsid-type structure.
- 35 28. Method according to one of the preceding claims, where one strand of the dsRNA is complementary to the primary or processed RNA transcript of the target gene.

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29. Method according to one of the preceding claims, where the cell is a vertebrate cell or a human cell.

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- 30. Method according to one of the preceding claims, where at least two dsRNAs which differ from each other are introduced into the cell, where at least segments of one strand of each dsRNA are complementary to in each case one of at least two different target genes.
 - 31. Method according to one of the preceding claims, where one of the target genes is the PKR gene.

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- 32. Medicament with at least one oligoribonucleotide with double-stranded structure (dsRNA) formed by two separate RNA single strands for inhibiting the expression of a given target gene, where one strand of the dsRNA has a region which is
- complementary to the target gene, characterized in that the complementary region has less than 25 successive nucleotide pairs.

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- 33. Medicament according to claim 32, where the dsRNA is enclosed by micellar structures, preferably by liposomes.
- 30 34. Medicament according to either of claims 32 or 33, where the dsRNA is enclosed by natural viral capsids or by chemically or enzymatically produced artificial capsids or structures derived therefrom.

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35. Medicament according to one of claims 32 to 34, where the target gene can be expressed in eukaryotic cells.

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36. Medicament according to one of claims 32 to 35, where the target gene is selected from the following group: oncogene, cytokin gene, Idprotein gene, development gene, prion gene.

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- 37. Medicament according to one of claims 32 to 36, where the target gene can be expressed in pathogenic organisms, preferably in plasmodia.
- 38. Medicament according to one of claims 32 to 37, where the target gene is part of a virus or viroid.
- 15 39. Medicament according to claim 38, where the virus is a virus or viroid which is pathogenic for humans.
- 40. Medicament according to claim 38, where the virus or viroid is a virus or viroid which is pathogenic for animals.
- 41. Medicament according to one of claims 32 to 40, where segments of the dsRNA are in double-stranded form.
 - 42. Medicament according to one of claims 32 to 40, where the ends of the dsRNA are modified in order to counteract degradation in the cell or dissociation into the single strands.
- 43. Medicament according to one of claims 32 to 42, where the cohesion of the double-stranded structure, which is caused by the complementary nucleotide pairs, is increased by at least one, preferably two, further chemical linkage(s).

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44. Medicament according to one of claims 32 to 43, where the chemical linkage is formed by a covalent or ionic bond, a hydrogen bond, hydrophobic interactions, preferably van-der-Waals or stacking interactions, or by metal-ion coordination.

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- 45. Medicament according to one of claims 32 to 44, where the chemical linkage is generated at at least one, preferably both, ends of the double-stranded structure.
- 46. Medicament according to one of claims 32 to 45, where the chemical linkage is formed by means of one or more compound groups, the compound groups preferably being poly(oxyphosphinicooxy-1,3-propanediol) and/or polyethylene glycol chains.
- 47. Medicament according to one of claims 32 to 46, where the chemical linkage is formed by purine analogs used in the double-stranded structure in place of purines.
- 48. Medicament according to one of claims 32 to 47, where the chemical linkage is formed by azabenzene units inserted into the double-stranded structure.
- 49. Medicament according to one of claims 32 to 48, where the chemical linkage is formed by branched nucleotide analogs used in the double-stranded structure in place of nucleotides.
- 50. Medicament according to one of claims 32 to 49, where at least one of the following groups is used for generating the chemical linkage: methylene blue; bifunctional groups, preferably bis(2-chloroethyl)amine; N-acetyl-N'-(p-glyoxylbenzoyl)-cystamine; 4-thiouracil; psoralene.

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51. Medicament according to one of claims 32 to 50, where the chemical linkage is formed by thiophosphoryl groups provided at the ends of the double-stranded structure.

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- 52. Medicament according to one of claims 32 to 51, where the chemical linkage are [sic] triple-helix bonds provided at the ends of the double-stranded structure.
- 53. Medicament according to one of claims 32 to 52, where at least one 2'-hydroxyl group of the nucleotides of the dsRNA in the double-stranded structure is replaced by a chemical group, preferably a 2'-amino or a 2'-methyl group.
- 54. Medicament according to one of claims 32 to 53, where at least one nucleotide in at least one strand of the double-stranded structure is a locked nucleotide with a sugar ring which is chemically modified, preferably by a 2'-0, 4'-C-methylene bridge.
- 25 55. Medicament according to one of claims 32 to 54, where the dsRNA is bound to, associated with or surrounded by, at least one viral coat protein which originates from a virus, is derived therefrom or has been prepared synthetically.
 - 56. Medicament according to one of claims 32 to 55, where the coat protein is derived from the polyomavirus.
- 35 57. Medicament according to one of claims 32 to 56, where the coat protein contains the polyomavirus virus protein 1 (VP1) and/or virus protein 2 (VP2).

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- 58. Medicament according to one of claims 32 to 57, where, when a capsid or capsid-type structure is formed from the coat protein, one side faces the interior of the capsid or capsid-type structure.
- 59. Medicament according to one of claims 32 to 58, where one strand of the dsRNA is complementary to the primary or processed RNA transcript of the target gene.
- 60. Medicament according to one of claims 32 to 59, where the cell is a vertebrate cell or a human cell.

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- 61. Medicament according to one of claims 32 to 60, where at least two dsRNAs which differ from each other are contained in the medicament, where at least segments of one strand of each dsRNA are complementary to in each case one of at least two different target genes.
- 62. Medicament according to claim 61, where one of the target genes is the PKR gene.

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63. Active ingredient with at least one oligoribonucleotide with double-stranded structure (dsRNA) formed by two separate RNA single strands for inhibiting the expression of a given target gene, where one strand of the dsRNA has a region which is complementary to the target gene, and where the target gene is part of a phytopathogenic virus or viroid,

characterized in that

the complementary region has less than 25 successive nucleotide pairs.

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- 64. Active ingredient according to claim 63, where the target gene can be expressed in eukaryotic cells.
- 65. Active ingredient according to claim 63 or 64, where segments of the dsRNA are in double-stranded form.
- 66. Active ingredient according to one of claims 63 to 65, where the ends of the dsRNA are modified in order to counteract degradation in the cell or dissociation into the single strands.
- 67. Active ingredient according to one of claims 63 to 66, where the cohesion of the double-stranded structure, which is caused by the complementary nucleotide pairs, is increased by at least one, preferably two, further chemical linkage(s).
- 68. Active ingredient according to one of claims 63 to 67, where the chemical linkage is formed by a covalent or ionic bond, a hydrogen bond, hydrophobic interactions, preferably van-der-Waals or stacking interactions, or by metal-ion coordination.
 - 69. Active ingredient according to one of claims 63 to 68, where the chemical linkage is generated at at least one, preferably both, ends of the double-stranded structure.
- 70. Active ingredient according to one of claims 63 to 69, where the chemical linkage is formed by means of one or more compound groups, the compound groups preferably being poly(oxyphosphinicooxy-1,3-propanediol) and/or polyethylene glycol chains.

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- 71. Active ingredient according to one of claims 63 to 70, where the chemical linkage is formed by purine analogs used in the double-stranded structure in place of purines.
- 72. Active ingredient according to one of claims 63 to 71, where the chemical linkage is formed by azabenzene units inserted into the double-stranded structure.

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- 73. Active ingredient according to one of claims 63 to 72, where the chemical linkage is formed by branched nucleotide analogs used in the double-stranded structure in place of nucleotides.
- 74. Active ingredient according to one of claims 63 to 73, where at least one of the following groups is used for generating the chemical linkage: methylene blue; bifunctional groups, preferably bis(2-chloroethyl)amine; N-acetyl-N'-(p-glyoxylbenzoyl)cystamine; 4-thiouracil; psoralene.
- 75. Active ingredient according to one of claims 63 to 74, where the chemical linkage is formed by thiophosphoryl groups provided at the ends of the double-stranded structure.
- 76. Active ingredient according to one of claims 63 to 75, where the chemical linkage are triple-helix bonds provided at the ends of the double-stranded structure.
- 77. Active ingredient according to one of claims 63 to 76, where at least one 2'-hydroxyl group of the nucleotides of the dsRNA in the double-stranded structure is replaced by a chemical group, preferably a 2'-amino or a 2'-methyl group.

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78. Active ingredient according to one of claims 63 to 77, where at least one nucleotides at least one strand of the double-stranded structure is a locked nucleotide with a sugar ring which is chemically modified, preferably by a 2'-0, 4'-C-methylene bridge.

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- 79. Active ingredient according to one of claims 63 to 78, where one strand of the dsRNA is complementary to the primary or processed RNA transcript of the target gene.
- 80. Active ingredient according to one of claims 63 to 79, where at least two dsRNAs which differ from each other are contained in the active ingredient, where at least segments of one strand of each dsRNA are complementary to in each case one of at least two different target genes.
- 20 81. Use of an oligoribonucleotide with double-stranded structure (dsRNA) formed by two separate RNA single strands or preparing a medicament or active ingredient for inhibiting the expression of a given target gene, where one strand of the dsRNA has a region which is complementary to the target gene,

characterized in that the complementary region has less than 25 successive nucleotide pairs.

- 82. Use according to claim 81, where the dsRNA is enclosed by micellar structures, preferably by liposomes.
- 35 83. Use according to either of claims 81 or 82, where the dsRNA is enclosed by natural viral capsids or by chemically or enzymatically produced artificial capsids or structures derived therefrom.

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- 84. Use according to one of claims 81 to 83, where the target gene can be expressed in eukaryotic cells.
- 5 85. Use according to one of claims 81 to 84, where the target gene is selected from the following group: oncogene, cytokin gene, Id-protein gene, development gene, prion gene.
- 10 86. Use according to one of claims 81 to 85, where the target gene can be expressed in pathogenic organisms, preferably in plasmodia.
- 87. Use according to one of claims 81 to 86, where the target gene is part of a virus or viroid.
 - 88. Use according to claim 87, where the virus is a virus or viroid which is pathogenic for humans.
- 20 89. Use according to claim 87, where the virus or viroid is a virus or viroid which is pathogenic for animals or phytopathogenic.
- 90. Use according to one of claims 81 to 89, where segments of the dsRNA are in double-stranded form.

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- 91. Use according to one of claims 81 to 90, where the ends of the dsRNA are modified in order to counteract degradation in the cell or dissociation into the single strands.
- 92. Use according to one of claims 81 to 91, where the cohesion of the double-stranded structure, which is caused by the complementary nucleotide pairs, is increased by at least one, preferably two, further chemical linkage(s).

***.**:

93. Use according to one of claims 81 to 92, where the chemical linkage is formed by a covalent or ionic bond, a hydrogen bond, hydrophobic interactions, preferably van-der-Waals or stacking interactions, or by metal-ion coordination.

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- 94. Use according to one of claims 81 to 93, where the chemical linkage is generated at at least one, preferably both, ends of the double-stranded structure.
- 95. Use according to one of claims 81 to 94, where the chemical linkage is formed by means of one or more compound groups, the compound groups preferably being poly(oxyphosphinicooxy-1,3-propanediol) and/or polyethylene glycol chains.
- 96. Use according to one of claims 81 to 95, where the chemical linkage is formed by purine analogs used in the double-stranded structure in place of purines.
- 97. Use according to one of claims 81 to 96, where the chemical linkage is formed by azabenzene units introduced into the double-stranded structure.
- 98. Use according to one of claims 81 to 97, where the chemical linkage is formed by branched nucleotide analogs used in the double-stranded structure in place of nucleotides.
- 99. Use according to one of claims 81 to 98, where at least one of the following groups is used for generating the chemical linkage: methylene blue; bifunctional groups, preferably bis(2-chloroethyl)amine; N-acetyl-N'-(p-glyoxylbenzoyl)-cystamine; 4-thiouracil; psoralene.

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100. Use according to one of claims 81 to 99, where the chemical linkage is formed by thiophosphoryl groups attached to the ends of the double-stranded structure.

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101. Use according to one of claims 81 to 100, where the chemical linkage at the ends of the double-stranded structure is formed by triple-helix bonds.

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102. Use according to one of claims 81 to 101, where at least one 2'-hydroxyl group of the nucleotides of the dsRNA in the double-stranded structure is replaced by a chemical group, preferably a 2'-amino or a 2'-methyl group.

103. Use according to one of claims 81 to 102, where at least one nucleotide in at least one strand of the double-stranded structure is a locked nucleotide with a sugar ring which is chemically modified,

preferably by a 2'-0, 4'-C-methylene bridge.

- 104. Use according to one of claims 81 to 103, where the dsRNA is bound to, associated with or surrounded by, at least one viral coat protein which originates from a virus, is derived therefrom or has been prepared synthetically.
- 105. Use according to one of claims 81 to 104, where the coat protein is derived from polyomavirus.
 - 106. Use according to one of claims 81 to 105, where the coat protein contains the polyomavirus virus protein 1 (VP1) and/or virus protein 2 (VP2).

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107. Use according to one of claims 81 to 106, where, when a capsid or capsid-type structure is formed

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from the coat protein, one side faces the interior of the capsid or capsid-type structure.

- 108. Use according to one of claims 81 to 107, where one strand of the dsRNA is complementary to the primary or processed RNA transcript of the target gene.
- 109. Use according to one of claims 81 to 108, where the cell is a vertebrate cell or a human cell.
- 110. Use according to one of claims 81 to 109, where at least two dsRNAs which differ from each other are used, where at least segments of one strand of each dsRNA are complementary to in each case one of at least two different target genes.
 - 111. Use according to claim 110, where one of the target genes is the PKR gene.

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112. Use according to one of claims 81 to 111, where the medicament is injectable into the bloodstream or into the interstitium of the organism to undergo therapy.

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- 113. Use according to one of claims 81 to 112, where the dsRNA is taken up into bacteria or microorganisms.
- 30 114. Use of a vector for coding at least one oligoribonucleotide with double-stranded structure (dsRNA) formed by two separate RNA single strands for preparing a medicament or active ingredient for inhibiting the expression of a given target gene, where one strand of the dsRNA has a region which is complementary to the target gene, characterized in that

the complementary region has less than 25 successive nucleotide pairs.

: .

- 115. Use according to claim 114, where the target gene can be expressed in eukaryotic cells.
- 116. Use according to claim 114 or 115, where the target gene is selected from the following group: oncogene, cytokin gene, Id-protein gene, development gene, prion gene.
 - 117. Use according to one of claims 114 to 116, where the target gene can be expressed in pathogenic organisms, preferably in plasmodia.
 - 118. Use according to one of claims 114 to 117, where the target gene is part of a virus or viroid.

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- 119. Use according to claim 118, where the virus is a virus or viroid which is pathogenic for humans.
 - 120. Use according to claim 118, where the virus or viroid is a virus or viroid which is pathogenic for animals or phytopathogenic.
 - 121. Use according to one of claims 114 to 120, where segments of the dsRNA are in double-stranded form.
- 122. Use according to one of claims 114 to 121, where one strand of the dsRNA is complementary to the primary or processed RNA transcript of the target gene.
- 123. Use according to one of claims 114 to 122, where the cell is a vertebrate cell or a human cell.
 - 124. Use according to one of claims 114 to 123, where at least two dsRNAs which differ from each other

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are used, where at least segments of one strand of each dsRNA are complementary to in each case one of at least two different target genes.

5 125. Use according to claim 125, where one of the target genes is the PKR gene.

Fetherstonhaugh & Co. Ottawa, Canada Patent Agents

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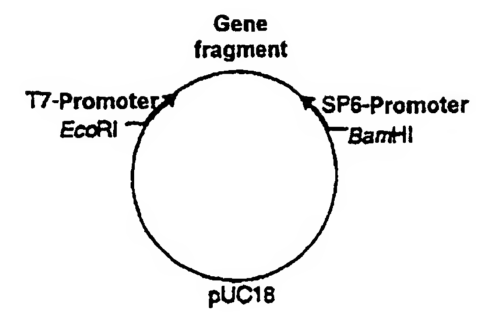


Fig. 1

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2/5%



Fig. 2

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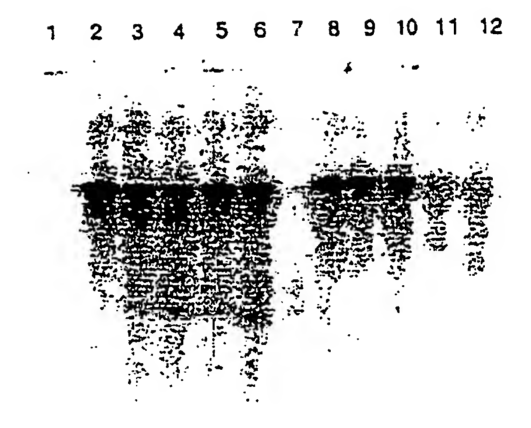
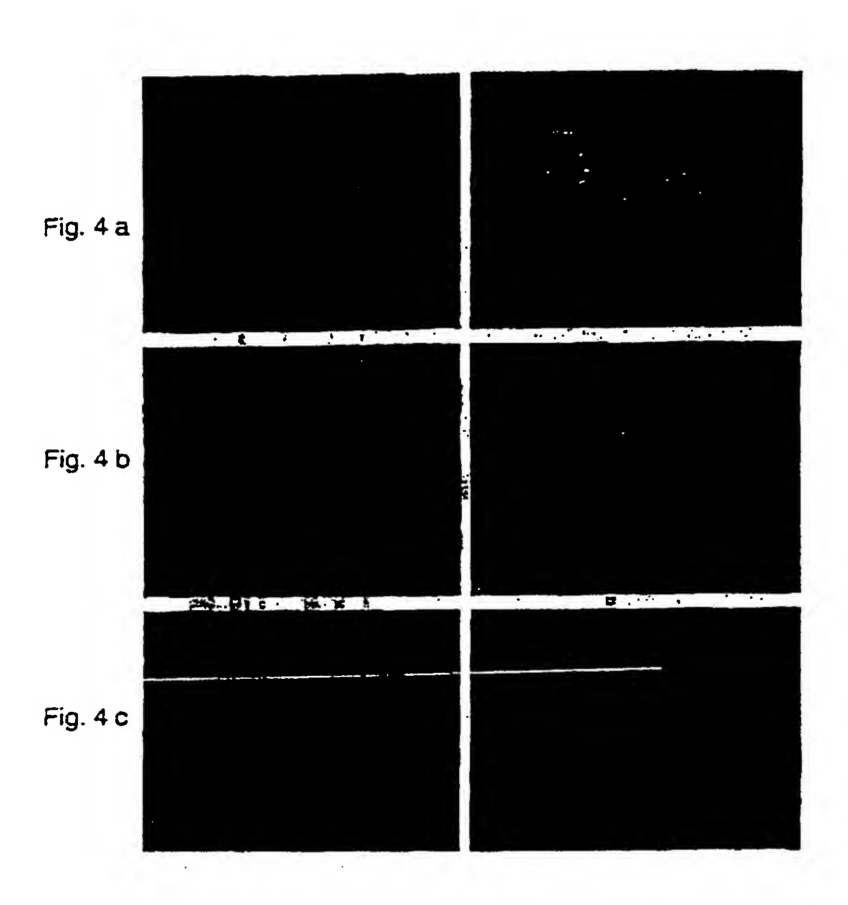
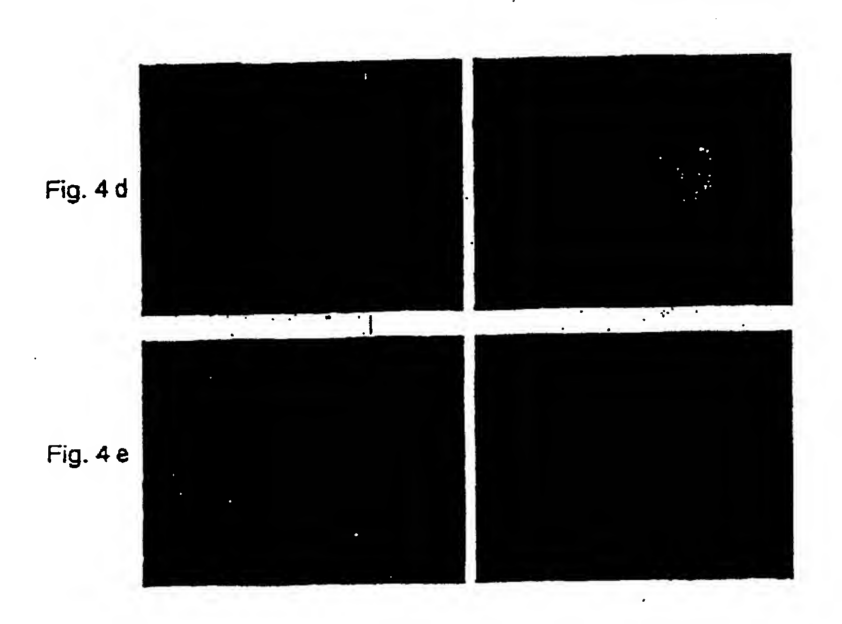


Fig. 3



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INTERNATIONAL SEARCH REPORT

Intl. Jonel Application No PCT/DE 00/00244

		PCT/DE 00	/00244
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	international Petent Classification (IPC) or to both national class SEARCHED	sification and IPC	
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	ENTS CONSIDERED TO BE RELEVANT		
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	abstract, page 11 lines 18-28 pages 12-13, page 15 line 22 bi pages 33 and 46, figures 1-6	s page 20 line 1,	1-35, 37-43, 45-72, 74-80, 82-108, 110-112
χ Funt	her documents are listed in the continuation of box C.	Patent family members are listed	in annex.
Special ca	stegaries of cited documents:		
"A" docume consider a surface which citation others	ent defining the general state of the art which is not dered to be of particular relevance document but published on or after the international	"I later document published after the into or priority date and not in conflict with cited to understand the principle or th invention. "X" document of particular relevance; the carnot be considered novel or canno involve an inventive step when the document of particular relevance; the cannot be considered to involve an indocument is combined with one or ments, such combination being obvior in the art. "&" document member of the same patent	the application but seary underlying the claimed invention to considered to ocument is taken alone claimed invention eventive step when the one other such document is a person skilled
Date of the	actual completion of the international search	Date of mailing of the international se	•
6	June 2000	20/06/2000	
Name and r	Trailing address of the ISA European Patent Office, P.8. 5818 Patentiasn 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 apo ni, Fax: (+31-70) 340-3016	Authorized officer Gore, V	

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hts. .tional Application No PCT/DE 00/00244

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	abstract, pages 6, 11-12, 15-17	112		
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	MADHUR K. ET AL.: "Antisense RNA: function and fate of duplex RNA in cells of higher eukaryotes." MICROBIOLOGY AND MOLECULAR BIOLOGY REVIEWS, vol. 62, December 1998 (1998-12), pages 1415-1434, XP000909741 * pages 1422-1423 and 1428 *	1-112		

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A. KLASSI IPK 7	FIZIERUNG DES ANMELDUNGSGEGENSTANDES C12N15/11 A61K31/713		
	ternationalen Patentidassifikation (IPK) oder nach der nationalen Klassif	Skation und der IPK	
	RCHIERTE GEBIETE		
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Withrend d	er Internationalen Recherche konautterte elektronische Datenbank (Nam	ne der Datenbank und evt. verwendete S	Suchbegriffe) .
C. ALS W	ESENTLICH ANGESEHENE UNTERLAGEN		
Kategorie*	Bezeichnung der Veröffentlichung, soweit erforderlich unter Angabe	der in Betracht kommenden Teile	Betr. Anspruch Nr.
Ϋ́	<pre># Zusammenfassung, Seite 11 Z.18-2 Seiten 12-13, Seite 15 Z.22 bis Seite</pre>		1-29, 32-34, 37-43, 45-66, 69-71, 74-80, 82-102, 105-108, 112, 1-35, 37-43,
	Z.1, Seiten 33 und 46, Abbildunger	1-6 * /	45-72, 74-80, 82-108, 110-112
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	6. Juni 2000 d Postanschrift der Internationalen Recherchenbehörde	20/00/2000 Bevollmächtigter Bediensteler	
	Europäisches Patentamt, P.B. 5818 Patentiaan 2 NL – 2280 HV Rijswijk Tel. (+31–70) 340–2040, Tx. 31 651 epo ni. Fax: (+31–70) 340–3016	Gore, V	

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Ints Jonales Aktenzeichen
PCT/DE 00/00244

		PCT/DE 00	00/00244	
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K	WO 98 05770 A (ROTHBARTH KARSTEN ; JOSWIG GABY (DE); WERNER DIETER (DE); SCHUBERT) 12. Februar 1998 (1998-02-12)		1-29, 32-34, 37-43, 45-66, 69-71, 74-80, 82-102, 105-108,	
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A ·	MADHUR K. ET AL.: "Antisense RNA: function and fate of duplex RNA in cells of higher eukaryotes." MICROBIOLOGY AND MOLECULAR BIOLOGY REVIEWS, Bd. 62, Dezember 1998 (1998-12), Seiten 1415-1434, XP000909741 Seiten 1422-1423 und 1428		1-112	

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Angaben zu Veröffentlichungen, die zur selben Patentiamilie gehören

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